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CONTROL OF NATURAL ZEOLITE PROPERTIES BY MECHANICAL ACTIVATION IN STIRRED MEDIA MILL

Due to the special characteristics of zeolites, they can be applied in a very wide range of industries, i.e. agricultural, environmental or water treatment purposes. Generally, high added value zeolite products are manufactured by micro- or nanogrinding. However, these processes require high energy input and cause significant wearing of the mill parts. Therefore, the optimization of zeolite grinding, as well as the control of its properties are of a great importance. In the present paper a Hungarian natural zeolite was mechanically activated in stirred media mill for various residence times in distilled water, meanwhile the particle size distribution and the grinding energy were measured. Additionally, on-line tube rheometer was used to study the rheology of the suspension during the grinding process. The particle interaction and the suspension aggregation stability were detected by zeta-potential measurements. Structural changes due to the mechanical activation process were monitored by FTIR. It was found that the material structure of the zeolite, as well as the rheological behaviour of the zeolite suspension and its aggregation stability had been altered due to the mechanical activation in the stirred media mill. It can be concluded that the zeolite product properties can be modified by mechanical activation in order to produce a high added value tailored material.

Keywords: zeolite, mechanical activation, zeta-potential, suspension rheology, FTIR

1. Introduction

Zeolites are natural crystalline aluminosilicates. Structurally zeolites are tectosilicates exhibiting an open three-dimensional structure containing cations needed to balance the electrostatic charge of the framework of silica and alumina tetrahedral. The collapse of the crystal structure can be desired in some cases since different strength of Bronsted and Lewis acid and base sites can thus be obtained [1,2]. This, in turn, can provide a better catalytic performance [3].

Ion-exchange properties of zeolites can be utilized in agriculture because of their large porosity and high cation-exchange capacity [4]. Additionally, they can be used as both carriers of nutrients and a medium to free nutrients [5]. Furthermore, zeolite incorporation in soil was found to increase crop yields and to promote nutrient uptake efficiency [6]. Other possible uses being investigated include application as a carrier of slow-release fertilizers, insecticides, fungicides, and herbicides, and as a trap for heavy metals in soils [7]. For groundwater remediation, permeable reactive barriers can be used [8] and zeolite can be applied as matrix material due to its high adsorption capacity and ion exchange capability.

Zolzaya et al [9] found that mechano-chemical activation in vibration mill improves cation exchange capacity of natural zeolites. Zeolite structure can be modified by acid or alkaline leaching to create mesopores and hence, to increase the surface area. At the same time Shumskaya et al. [10] suggested that mechano-chemical activation of zeolite improve its reactivity due to the ruptures of the Al-O-Si bonds in zeolite. This wide range of possible applications elevates zeolite among other strategic minerals [11] in Hungary.

Nanosize natural zeolites can be manufactured by micro- or nanogrinding, for example in stirred media mill, which is a high energy density mill. The movement of the grinding medium is carried out by the rotation of the concentrically or eccentrically placed rotor. As a result, the grinding balls colliding with each other are pressed to the mill liners; and collide with the rotor. The particles of the feed material are fractured by impact, pressure, collide and friction stresses [12].

Stirred media mills can be used for various aims, such as mechano-chemical activation of cementitious materials [13]. It can be used also for the enhancement of mass transfer in bio- and chemical processes and ultrafine liberation of semiconductors from glass substrate [14]. These mills can be used in dry [15] or wet [16-20] mode.

The main aim of this research was to investigate the mechanical activation of natural zeolite in a high energy density mill in order to produce submicron final product with narrow size distribution and desired particle interaction properties (stability). Material structure, rheological properties and aggregation stability of the zeolite suspension were examined.

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2. Materials and methods

2.1. Material

Natural zeolite with material density of 2200 kg/m³, originated from Rátka deposit, Hungary was used in this study. Its main mineral components were identified as clinoptilolite, orthoclase, cristobalite and quartz by XRD measurement using a Bruker D8 apparatus. Mineral composition is presented in Table 1. Cristobalite and quartz are usually observed in clinoptilolite rich zeolitic tuffs [21].

TABLE 1 Mineral composition of natural zeolite used in the experiments (in w/w%, dry basis)

Phase name	Wt% Rietveld
Clinoptilolite	38.2
Quartz	9.6
Cristobalite low	17.3
Orthoclase	11.3
Sanidine Na0.16	2.8
Illite 2M1	7.7
Cristobalite high	0.8
Mordenite	1.3
Amorphous	11.0
Sum	100.0

The taken natural zeolite sample was preliminary crushed, ground and classified; the maximal particle size of the product was less than 116 μ m. The volume ratio particle size distribution shows bimodal nature (Fig. 3).

2.2. Methods

2.2.1. Mechanical activation by grinding

Grinding experiments were carried out in a stirred media mill (Netzsch MiniCer, Netzsch GmbH, Germany) in continuous closed circuit mode. The liners of the grinding chamber and the stirrer rotors were made of high wear resistant ZrO (zirconia). The revolution number of the stirrer was 3000 min⁻¹ (9.3 ms⁻¹ circumferential tip speed) during the experiment. Mill volume in circuit mode is 500 ml. With the tube rheometer the volume can reach 805 ml. The grinding media size range was 500-600 μ m ceramic beads made of yttrium stabilized zirconium oxide (ZY Premium, Sigmund and Lindner, Germany) were used. The filling ratio of grinding media in the milling chamber was 70 v/v %.

Samples were taken after 5, 10, 20, 30, 60, 120, 240, 360 and 480 min experiment times then analysed using laser particle size analyser (LPSA). Samples were taken from the outflow of the pipe in which the suspension re-enters into the stirrer vessel. The volumetric flow rate was $2.5e-05 \text{ m}^3/\text{s}$ in case of 190 rpm rotation speed of the pump. The power draw of the mill was measured online by the Netzsch measuring system and data were registered in the computer. In this way the specific grinding energy could be measured.

2.2.2. Particle size distribution

The particle size distribution (PSD) of the raw material and the ground samples was determined by a Horiba LA-950V2 laser particle size analyser (LPSA) in distilled water, using ultrasonic dispersing for 1 min. Furthermore, the "outer" (geometric) specific surface area (SSA) of the ground samples was calculated by the same LPSA using shape factor 1.0 (Heywood factor).

2.2.3. Rheology

In a previous research Faitli et al. [22] introduced an online rheological testing equipment, in which the Netzsch stirred media mill was modified and a supplementary on-line tube rheometer was developed and installed. The laminar pipe flow is a fundamental phenomenon and therefore the tube rheometer is a suitable instrument to measure non-Newtonian rheological behaviour.

The original peristaltic pump of the Netzsch MiniCer was used as the main pump to drive the suspension through the measuring pipe sections. Two measuring pipe sections made from stainless steel were connected serially ($D_1 = 10$ mm, $L_1 = 1.6$ m, $D_2 = 9$ mm, $L_2 = 1.6$ m). If there are two measured pressure loss – flow rate points for a given operational condition, a two parameters non-Newtonian rheological model can be fit. Of-course, the flow rate can be changed and many points of the pseudo shear curve can be measured. The pressure loss of the test pipe sections were measured by no dead space gauge pressure transducers ($\Delta p_1 = p_1 - p_2$; $\Delta p_2 = p_3 - p_4$). For the tube rheometer an independent computer data acquisition system was developed and signals of the pressure transducers No. 1-4 were measured by this auxiliary system with 100 Pa sensitivity.

The method of evaluation of Bingham plastic tube rheometer data and the validation of tube rheometer data by a rotational rheometer can be found in a previous work [23].

2.2.4. FTIR

To follow the structural changes due to the mechanical activation process of zeolite a JASCO FT-IR 4200 type Fourier Transformed Infrared Spectrometer in Reflection mode with diamond ATR PRO470-H condition was used. Three spectra were made from each part-samples and every sample was measured with 3 part-samples. One spectrum averaged of 64 FTIR measurements with 4 cm⁻¹ resolution number. The incident angle was 45°, TGS detector was used and scanning speed was 2 mm/sec.

The ζ -potential measurements were carried out by a Zeta-Pals device from Brookhaven Instrument Corporation (BIC). The equipment determines zeta-potential using Phase Analysis Light Scattering (PALS), which is an extension of laser electrophoretic light scattering (ELS). ζ -potential was calculated from the electrophoretic mobility using the Smoluchowsky model. To ensure the reproducibility of the measurements, one measurement involved at least 10 several runs. The number of cycles in a run was 20. Zeta potential measurements were carried out within the whole continuous grinding experiment with the freshly taken, as well as with the samples after several hours long steady conditioning. Before the measurements the samples were diluted with de-ionised water and stirred with a magnetic stirrer.

3. Results and discussion

3.1. Mechanical activation by grinding

The particle size distribution curves of the ground samples are shown in Fig. 1. Rapid decrease in zeolite fineness was observed already after 30 min, followed by a section where particles in a narrower size range were generated. The median and x_{80} cumulative particle size were 0.21 µm and 0.29 µm after 120 min residence time respectively. The lowest median and x_{80} particle size were reached after 360 min residence time, found to be 0.15 µm and 0.22 µm respectively.



Fig. 1. Particle size distributions of the raw zeolite (feed) and ground products

Fig. 2 represents the x_{80} and x_{95} cumulative passing particle size changing during grinding. The x_{80} characteristic particle size decreased continuously during grinding from 26 μ m down

to 0.33 μ m. Meanwhile, the x₉₅ characteristic particle size decreased from the initial 57 μ m down to 0.55 μ m at 90 minutes grinding time, then it was increased up to 3.0 μ m at 240 minutes grinding time then it was nearly constant.



Fig. 2. The x₈₀ and x₉₅ particle size of zeolite during grinding

In Fig. 3 it can be seen that bimodal nature of the histogram of the raw zeolite remained characterising in the first stage of grinding, but the mode values were significantly shifted to the submicronic size range. It can also be seen clearly that not only the surface attrition, but breakage of the particles occurred in the grinding process.



Fig. 3. Histograms of raw zeolite and ground products

The calculated specific surface area as a function of the specific energy input (Fig. 4) initially increases near linearly



Fig. 4. The calculated specific surface area as a function of the specific energy

(until 1000 kJ/kg, SSA = 12 m²/g), this interval is generally referred as the Rittinger section of grinding kinetics. This is followed by a fluctuating curve shape indicating the aggregation or agglomeration of the primary zeolite particles and the generation of secondary ones. However, the maximum SSA, as high as 18 m²/g, was reached at 17,000 kJ/kg specific energy after 360 min grinding time.

3.2. Rheology

Fig. 5 shows the apparent viscosity of the suspension as a function of residence time in case of the tube rheometer test using one measuring tube. As it is well known the apparent vis-



Fig. 5. Apparent viscosity of the ground suspension as function of time

cosity is not a rheological material parameter like the absolute viscosity of Newtonian fluids. The magnitude of the apparent viscosity is proportional with the inner resistance of the fluid against the movement. The inner viscous resistance of the suspension increased initially, and at 35 minutes it reached a maximal peak value of 0.034 Pas. After this peak, it started to decrease and after 300 minutes stagnated until t = 480 minutes.

After 480 minutes of continuous grinding the mill was stopped and an independent tube rheometer test was performed. During this rheological test the revolution number of the peristaltic pump was systematically varied, so different flow rates were set in the measuring tube section. The measured pseudo shear curve is shown in Fig. 5.



Fig. 6. The measured pseudo shear curve after 480 min long grinding

Fig. 6 shows that there are measured shearing points only in the 270-400 1/s pseudo shear rate range. It is not possible to measure at zero or near zero flow velocity (shear rate) with the tube rheometer. Based on Fig. 6 the Bingham plastic linear and the Power Law exponential (shear thinning) rheological models could be applicable. The linear model fits slightly better because the coefficient of determination is a little bit higher; therefore the suspension can be considered as being a Bingham plastic fluid. The yield stress is $\tau_0 = 3 \cdot b/4 = 1.6$ Pa and the plastic viscosity is $\eta = a = 14$ mPa \cdot s.

3.3. FTIR

FTIR spectra is suitable to acquire information about the structure, structure change and the cation substitution (Si^{4+} by AI^{3+}) in the tetrahedral sites of zeolite minerals [24-28]. The results of FTIR measurements are shown in Fig. 7 where the characteristic bonds [29] are marked and the results are summarized in Table 2.

Sample	Feed	30 min	60 min	120 min	240 min	360 min	480 min
Symmetric stretching vibration (Si–O–Si and Al–O–Si)	594	590	594	595	592	592	553, 565, 591
Symmetric stretching vibration (Si–O–Si and Al–O–Si)	729	721	720	721	716	714	722
Symmetric stretching vibration of Si-O-Si(Al)	795	796	795	795	794	793	792
Asymmetric stretching vibration (Si(Al-O)	1011	1011	1018	1014	1015	1021	1012, 1093
Bending vibrations (HOH)	1631	1633	1630	1631	1643	1639	1636, 1650
Stretching vibration (-OH, HOH)	3407	3417	3432	3421	3404	3401	3269. 3365

The exact position (peak wavenumber) of bonds observed in FTIR spectra for raw and ground zeolite

Bands due to the Si-O-Si and Si-O-Al vibrations occur in the region 1200-950 cm⁻¹, where the strongest vibration band appears at 1010 cm⁻¹. It was found in all zeolites due to the internal tetrahedron vibrations and is assigned to a T-O stretching mode [30]. This band (at 1010 cm⁻¹) is significant for the estimation of the aluminium content in the crystalline framework. It depends on the framework nSi/nAl ratio and is considered to be determination for the Al atoms per unit. It shifts to a lower wavenumber with increasing number of Al atoms in the framework tetrahedral sites [31]. Moreover, substitution of Al³⁺ for Si⁴⁺ decreases T-O-T angle, and the resulting frequency is lower, due to the weaker character and increased length of the Al-O bond, the force constant for the mode in an Al-Si-O bond is lower than the corresponding mode is a Si-O-Si linkage. Thus, it is expected that the more Al³⁺ atoms are extracted from the framework, the more this band will shift to higher wave number values in the FTIR spectra [24]. At the activated sample after 60 min of grinding it shifts to 1018 cm⁻¹ and after 360 min to 1021 cm⁻¹.

In all FTIR spectra, the bands are quite broad, which is owing to the fact that all samples consist of a variety of crystalline phases and amorphous mass [32]. A weak band at 795 cm⁻¹ appears in all spectra and can be assigned to quartz or amorphous SiO₂ stretching bond vibrations. The vibrations in the region 1600-3700 cm⁻¹ can be assigned to the presence of zeolite water. The bands are attributed to the hydrogen bonding of the water molecule to surface oxygen (3407 cm⁻¹) and to the bending mode of water (1631 cm⁻¹) [24].

In the region 600-800 cm⁻¹ (590 and 720 cm⁻¹), bands attributed to exchangeable cations may occur, but bands due to pseudocrystallic vibrations may be confusing [32,33] These bands are clearly lower intensity in the FTIR spectra of 360 and 480 min samples – as compared to feed sample – as these samples could suffer crystallinity loss. These vibrations are insensitive to the nature of the channel cations, as well as to the Si/Al ratio.

New bonds appeared in case of the 480 min ground sample as symmetric (Si-O-Si(Al)) at 553, 565 cm⁻¹, asymmetric stretching vibrations (Si(Al-O)) at 1093 cm⁻¹, bending vibrations (HOH) at 1650 cm⁻¹ and stretching vibration (-OH, HOH) at 3269 and 3365 cm⁻¹. The shoulder at 1093 cm⁻¹ is due to the presence of cristobalite and smectite, with the possible contribution of amorphous silica [24].

Peak shift also can be observed. The most significant shifts are in case of symmetric stretching vibrations (Si-O-Si(Al)) and



Fig. 7. FTIR spectra of zeolite samples during grinding

stretching vibration (-OH, HOH) at $3300-3400 \text{ cm}^{-1}$. Based on these results it can be stated that the material structure of the zeo-lite sample had been changed due to the mechanical activation.

Fig. 8 represents the main peaks shifted during the grinding. The absolute differences between the ground samples peaks' wavenumber and the feed material peak's wavenumber are plotted as a function of grinding time. It can be observed that the HOH bending vibration at ~1630 cm⁻¹ wavenumber peak changed in the opposite way as compared with the stretching vibration –OH, HOH at ~3400 cm⁻¹. Asymmetric stretching vibration (Si(Al-O)) at 1011 cm⁻¹ also shifted to a higher wave number values especially at 30 and 360 min resulted from Al³⁺ atoms are extraction from the framework.

Fig. 9 shows intensity change of FTIR peaks compared with the initial state. The intensity change is the most significant at the asymmetric stretching vibration (Si(Al-O) at ~1010 cm⁻¹ and the symmetric stretching vibration (Si–O–Si and Al–O–Si) at 590 cm⁻¹ during grinding. They show three intensity maxima at 30,120 and 360 minutes grinding time as compared to the feed material. The highest maximum is at 360 minutes in both cases,



Fig. 8. FTIR peak shifting during grinding at 1630 and 3400 $\rm cm^{-1}$ wavenumber

16 and 8%T. The first two intensity maxima coincide with the i.e.p.-s, while the third one with the minimum of the $\xi = \xi(t)$ curve obtained for the freshly ground sample. Nevertheless, the first intensity maximum, as well as the first i.e.p. coincides with the peak point of the apparent viscosity. Furthermore, X-ray diffraction measurement for the precise determination of the mineralogical phase changes will be carried out in near future.

3.4. Zeta-potential

Zeta-potential measurements were carried out aiming at the revelation of possible changes in the surface charge and sign, as well as in the aggregation stability of zeolite during the 480 min long continuous micro-grinding [14,34-37].

The results are shown in Fig. 10. It can be seen in case of the freshly ground samples (green curve) that zeta-potential of the zeolite in the first 60 min of retention time of grinding follows a minimum-curve with the lowest ζ -potential of -2.35 mV. It reaches the highest value of 5.1 mV at t = 60 min. During the first 60 min there is no aggregation stability in the system, the particle-particle aggregation is the most likely here. During the next 300 min the significant drop in ζ -potential was observed, down to -32.64 mV with the isoelectric point, the i.e.p. at t = 125 min. In the grinding time period between 150 and 360 min the system is aggregatively stable due to the significant electrostatic repulsion of the negatively charged zeolite particles. In the last tested time period the ζ -potential increases again up to -11.7 mV. The curve trend shows that the aggregation stability might turn into instability during possible further grinding. The initial pH-value was as high as 8.34, and it dropped down to pH = 7.87 during the grinding.



Fig. 9. FTIR peak intensity changing during grinding at 1010 and 590 $\rm cm^{-1}$ wavenumber



Fig. 10. Zeta-potential of zeolite during the continuous micro-grinding

The ζ -potential curve obtained for the several hours long steady conditioning looks like the green curve would be shifted to the left with the first minimum at t = 180 min (ζ -potential as low as -38.11 mV). Although, it has a second minimum as well (-29.67 mV at 420 min) and crosses the green curve twice until reaches it.

It can be revealed and concluded, that at the early kinetic stage of the grinding of zeolite mostly the =Si=O bonds suffer breakage, while at continuous grinding longer than 125 min, predominantly the =Al-O- bonds are breaking providing a high negative surface charge, i.e. the aggregation stability. The even longer continuous grinding leads to the surplus of surface en-

ergy in such extend that the system tends to compensate it, i.e. zeta-potential increases and the aggregation might take place.

The zeolite samples after different grinding time were then steady conditioned for several, from 2.5 up to 15 hours. The zetra-potential curve of the steady conditioned samples is plotted against the real grinding time and shown in black. It can be seen that there is a shift of zeta-curve to the left below t = 240 min. It is very likely that some kind of restructuring of the microground structure of zeolite takes place in this interval. So that, the obtained structure alteration of the zeolite is a reversible one in this interval. At higher grinding time the difference between the two zeta-curves is not significant.

The correlation between the zeta-potential and the apparent viscosity of the tested zeolite system is shown in Fig. 11.



Fig. 11. Zeta potential as a function of apparent viscosity

In the tested interval increasing from the aggregation stability due to the negative surface charge, up to the aggregation non-stability (-5...+5 mV) of the ξ -potential of freshly ground samples, the apparent viscosity grows.

Zeolite at the examined weak alkaline conditions (pH = 7.87...8.34) has slightly positive surface charge. However, after 60 minutes of grinding the positive charge became slightly higher. As a consequence of the grinding the newer and newer surface appears, as well as the rupture of chemical bonds occurs. In the FTIR spectra of the freshly ground sample at 60 min the band at 1011 cm⁻¹ shifted to the left (higher wave number values), in terms of literature due to the decrease in number of Al³⁺ atoms in the framework tetrahedral sites [24,30,31], so that Si⁴⁺ become potential-determining ions.

The next point, where more Al^{3+} atoms are supposed to be broken off from the framework, in accordance with the FTIR spectra of the 360 min sample. At this point the specific surface area has maximum (18 m²/g) and the x₈₀ characteristic particle size has minimum (0.22 µm) value. It follows that the most surface during grinding formed at this grinding time. In this point the zeta-potential has minimum value which indicates negatively charge ions and groups such as OH^- , O^{2-} , $Si-O^{2-}$ closely approaches the surface and adsorb on it. As a result of further grinding, the aggregation or agglomeration can occur as well, this is proved by the increasing of specific surface area and decreasing of zeta-potential and characteristic particle size (x_{80} and x_{95}).

The relationship between the apparent viscosity and the zeta-potential can be observed quantitatively, since both phenomena originate from the interaction between solid-solid and solid-liquid particles. Around the i.e.p. of zeta-potential the apparent viscosity has maximum value (Fig. 11). Because of aggregation, the apparent viscosity should increase.

4. Conclusions

Nano zeolite was successfully produced using Netzsch MiniCer stirred media mill. The mode size of 200 nm and the highest geometric specific surface area of $18 \text{ m}^2/\text{g}$ was achieved after 360 min continuous grinding. It was experimentally proved that the material structure of zeolite, the rheological property and the aggregation stability of the zeolite suspension have been altered due to the mechanical activation in the stirred media mill. The interrelations between the structural changes, the zetapotential, as well as the rheological behaviour were revealed. The obtained results are very promising from the point of view of micro- and nanogrinding of tailored products.

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